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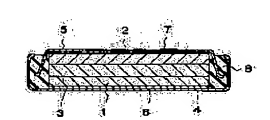
KOJIMA TOSHIAKI

(54) LITHIUM SECONDARY BATTERY

(57)Abstract:

PROBLEM TO BE SOLVED: To provide a lithium secondary battery having high capacity, high energy density, low irreversible capacity, and excellent charge and discharge characteristic, even in combining with a carbon material, and high temperature.

SOLUTION: This lithium secondary battery using a carbon material as a main constituent of a negative electrode active material uses Li1-z [Mn2-x-yMxLiyO4] as a main constituent of a positive electrode active material (M is an element except Mn and Li, then (x) and (y) represent an amount of substitution of manganese, and they meet the following inequality $0 \le x+y \le 1$. Further, (z) represents an amount of reversibly usable lithium, and it meets the following inequality, $-1 \le z \le 1$).



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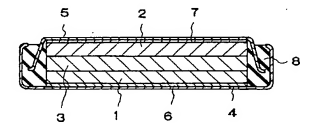
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(54) 【発明の名称】 リチウム二次電池

(57)【要約】

(目的) 炭素材料との組合せや高温においても高容量、高エネルギー密度で、不可逆容量の少ない充放電サイクル特性の優れたリチウム二次電池を提供することを目的とする。

【構成】 負極活物質の主構成物質として炭素材料を用いるリチウム二次電池において、正極活物質の主構成物質にし i_{1-2} [Mn_{2-x-y} M_x Li_v O_x] (MidMn Li_v U_x U_y $U_$



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【特許請求の範囲】

【請求項1】 負極活物質の主構成物質として炭素材料を用いるリチウム二次電池において、正極活物質の主構成物質にLi_{1-z} [Mn_{2-x-v} M_x Li_v O₄] (Mは MnとLi以外の元素で、x,yはマンガンの置換量を示し、 $0 < x + y \le 1$ である。また、zは可逆的に使用可能なリチウム量を示し、 $-1 \le z \le 1$ である。)を用いることを特徴とするリチウム二次電池。

【請求項2】 前記正極活物質の主構成物質が、Li $_{1-x}$ [Mn_{2-x-v} $Ca_xLi_vO_v$] (x,yはマンガンの置換量を示し、 $0 < x + y \le 1$ である。また、z は可逆的に使用可能なリチウム量を示し、 $-1 \le z \le 1$ である。)を用いることを特徴とする請求項1記載のリチウム二次電池。

【請求項3】 前記正極活物質の主構成物質が、Li1-x [Mn_{x-x-} , $Cr_xLi_vO_v$](x, yはマンガンの置換量を示し、0 < x + y ≤ 1 である。また、z は可逆的に使用可能なリチウム量を示し、-1 $\leq z$ ≤ 1 である。)を用いることを特徴とする請求項1記載のリチウム二次電池。

【請求項4】 前記正極活物質の主構成物質が、Li $_{1-z}$ $[Mn_{z-x-v}$ $In_xLi_vO_t$] (x,yはマンガンの置換量を示し、 $0 < x + y \le 1$ である。また、z は可逆的に使用可能なリチウム量を示し、 $-1 \le z \le 1$ である。)を用いることを特徴とする請求項1記載のリチウム二次電池。

【請求項5 】 前記正極活物質の主構成物質が、 L_1 1 = 1 = 1 1

【請求項6】 前記正極活物質の主構成物質が、Li1-1、 $[Mn_{1-x-}, Mg_*Li_*, O_*]$ (x, yはマンガンの置換量を示し、0 < x $+ y \le 1$ である。また、z は可逆的に使用可能なリチウム量を示し、 $-1 \le z \le 1$ である。)を用いることを特徴とする請求項1 記載のリチウム二次電池。

【請求項7】 前記負極活物質である炭素材料が、X線回折法による面間隔(d002)が3.354~3.369Åで、C軸方向の結晶の大きさ(Lc)が200Å以上であることを特徴とする請求項1記載のリチウム二次電池。

【発明の詳細な説明】

[0001]

【発明の属する技術分野】本発明はリチウム二次電池に係り、放電容量、出力密度が大であって、特に高温でのサイクル特性に優れた安全性の高いリチウム二次電池用 正極及び負極に関する。

[0002]

【従来の技術】現在、4Vを特徴とするリチウム二次電 池の正極活物質として、LiCoO,、LiNiO,等 のα-NaFeO、構造を有する含リチウム酸化物、あ るいはLiMn、O、等のスピネル構造を有する含リチ ウム酸化物等が従来より用いられている。なかでもスピ ネル構造を有するLiMn、O、は、低コストかつ安全 性に優れた正極活物質である。一方、負極活物質として は、従来よりリチウム金属及びリチウム合金が用いられ てきたが、これらの電池は、樹枝状リチウムの析出(デ 10 ンドライト) による正負両極の短絡やサイクル寿命が短 く、そのためその劣化分を補償すべく電池容量の3倍当 量のリチウムが必要であり、エネルギー密度が低いとい う欠点があった。最近ではこれらの問題点を解決するた め炭素材料を負極に用いる研究が活発である。この種の 負極、特に黒鉛化の進んだグラファイトを用いる場合、 例えば正極にマンガン酸リチウムを用いると、電池電圧 がフラットなものになり、単電池使用の携帯機器に用い る場合容量面で優位性がある。しかしながら、負極グラ ファイトを、正極にLiMn、〇、を用いて充放電サイ 20 クルを行うと、充電時のグラファイトのドープ電圧がサ イクルを重ねるとOV付近となり、リチウムの析出との 競争反応となってしまう。そのため、サイクル劣化が大 きい要因の一つとなっていた。また、このマンガン酸リ チウムはその正極活物質自身のサイクル特性が優れない ことから、特開平4-233161号公報や特開平5-21067号公報や特開平6-187993号公報にお いてはスピネル構造のマンガン酸リチウムのマンガンの 一部をマンガン以外の元素で置換することが報告されて いる。しかしながら、これらの材料においても髙温で放 置しておくとマンガンが電解液に溶出することから、正 30 極の充放電容量が減少し、サイクル劣化することが分か った。

[0003]

【発明が解決しようとする課題】前述した如く、炭素材料を負極として用いた場合、マンガン酸リチウムとの組合せによって炭素の界面抵抗の増加という問題がある。また、正極活物質自身高温での充放電によって充放電容量が低下するという問題もある。本発明は、この問題点を解決するため、正極活物質における主構成物質に、Li、[Mnz...、Mx Li、O 、](x、yはマンガンの置換量を示し、 $0 < x + y \le 1$ である。また、zは可逆的に使用可能なリチウム量を示し、 $-1 \le z \le 1$ である。)を用いることにより、炭素材料との組合せや高温においても高容量、高エネルギー密度で、不可逆容量の少ない充放電サイクル特性の優れたリチウム二次電池を提供することを目的とする。

[0004]

【課題を解決するための手段】負極活物質として炭素を 考えた場合、炭素材料へのリチウムの吸蔵、放出(イン 50 ターカレーション、デインターカレーション)が主に起

こる反応だが、その反応を支配する因子の一つとして、 電解液と炭素表面の間に生じる被膜状態が関与している ことがわかった。例えば、リチウム金属を負極活物質に した場合で代表されるように、緻密でイオン導伝性の高 い被膜はその電池特性も優れており、逆に厚くイオン伝 導性の低い被膜はレート特性や、サイクル特性が悪いこ とが知られている。その場合、前者は炭酸リチウムや酸 化リチウム等の被膜であり、後者はフッ化リチウム等の 被膜であることが報告されている。これと同じことが炭 素表面に生じる被膜についても考えられる。つまり、炭 10 素材料の界面抵抗を増大させる要因の一つとして、炭素 材料の表面にフッ化リチウム等のイオン伝導度の低い被 膜の形成があげられる。このフッ化リチウムの被膜が形 成される過程において、正極材料等から持ち込まれた水 により電解質が分解して生成するフッ酸が関与している ことが考えられる。本発明者らは、この問題点を解決す るため種々検討した結果、スピネル構造を有するマンガ ン酸リチウムにおいて、マンガンの一部をリチウムと、 カルシウム、クロム、インジウム、テルビウム、マグネ シウム等のマンガン以外の元素で置換することにより、 電池内部で生成するフッ酸を抑制することができること を見い出した。また、この正極材料は髙温保存による電 解液へのマンガンの溶出も抑制することが分かった。

【0005】正極活物質の主構成物質であるLi 1-2 [Mn2-x-y Mx Li, O,] において、MはMn とLi以外の元素で、Mnと置換しうる元素が好まし い。例えば、Be, B, C, Si, P, Sc, Cu, Z n, Ga, Ge, As, Se, Sr, Mo, Pd, A g, Cd, In, Sn, Sb, Te, Ba, Ta, W. Pb, Bi, Co, Fe, Cr, Ni, Ti, Zr, N b, Y, Al, Na, K, Mg, Ca, Cs, La, C e, Nd, Sm, Eu, Tb等が挙げられる。これらの 中でもCr、Ca、In, Tb, Mgはその効果が顕著 であったため最も好ましい。ここで、Mnの置換量を示 すx, yは、0 < x + y ≤ 1 である。ただし、Mnの置 換量x+yを増やすと、可逆的に使用可能なリチウム量 を示すzが減少するため、0<x+y≤0.2の範囲で 置換することが好まし。可逆的に使用可能なリチウム量 を示すzは、Mnの置換量によっても異なってくるが一

【0006】正極活物質の主構成物質であるLi 1-, [Mn_{2-x-y} M_x Li, O,] において、Mnの一 部をリチウムや異種元素で置換する場合、焼成原料にあ らかじめ置換する元素を添加する方法や、LiMn、O 、を焼成した後にイオン交換等により異種元素を置換す る方法等が挙げられるが、これらに限定されるものでは ない。

【0007】負極活物質に用いる炭素材料は、リチウム を吸蔵、放出可能な炭素材料であればよく、特にX線回

9Åで、C軸方向の結晶の大きさ(Lc)が200Å以 上である炭素粒子は、髙容量が得られるため好ましい。 【0008】本発明に用いる正極、負極活物質は、平均 粒子サイズ100μm以下であることが望ましい。所定 の形状を得る上で、粉体を得るためには粉砕機や分級機 が用いられる。例えば、乳鉢、ボールミル、サンドミ ル、振動ボールミル、遊星ボールミル、ジェットミル、 カウンタージェットミル、旋回気流型ジェットミル、熱 プラズマや篩等が用いられる。粉砕時には水、あるいは ヘキサン等の有機溶剤を共存させた湿式粉砕を用いると ともできる。分級方法としては、特に限定はなく、篩や 風力分級機などが乾式、湿式ともに必要に応じて用いら れる。

【0009】本発明に併せて用いることができる負極材 料としては、リチウム金属、リチウム合金などや、カル コゲン化合物、メチルリチウム等のリチウムを含有する 有機化合物等が挙げられる。また、リチウム金属やリチ ウム合金、リチウムを含有する有機化合物を併用すると とによって、本発明に用いる炭素材料にあらかじめリチ ウムを挿入することも可能である。

【0010】本発明の正極、負極活物質を用いる場合、 電極合剤として導電剤や結着剤やフィラー等を添加する ことができる。導電剤としては、電池性能に悪影響を及 ほさない電子伝導性材料であれば何でも良い。通常、天 然黒鉛(鱗状黒鉛、鱗片状黒鉛、土状黒鉛など)、人造 **里鉛、カーボンブラック、アセチレンブラック、ケッチ** ェンブラック、カーボンウイスカー、炭素繊維や金属 (銅、ニッケル、アルミニウム、銀、金など)粉、金属 繊維、導電性セラミックス材料等の導電性材料を1種ま たはそれらの混合物として含ませることができる。これ らの中で、アセチレンブラックとケッチェンブラックの 併用が望ましい。その添加量は1~50重量%が好まし く、特に2~30重量%が好ましい。

【0011】本発明の正極、負極活物質を用いる場合、 その粉体の少なくとも表面層部分を主活物質以外のもの で修飾することも可能である。例えば、金、銀、カーボ ン、ニッケル、銅等の電子伝導性のよい物質や、炭酸リ チウム、ホウ素ガラス、固体電解質等のイオン伝導性の よい物質をメッキ、焼結、メカノフュージョン、蒸着等 40 の技術を応用してコートすることが挙げられる。

【0012】結着剤としては、通常、テトラフルオロエ チレン、ポリフッ化ビニリデン、ポリエチレン、ポリプ ロピレン、エチレン・プロピレンジエンターポリマー (EPDM)、スルホン化EPDM、スチレンブタジエ ンゴム(SBR)、フッ素ゴム、カルボキシメチルセル ロース等といった熱可塑性樹脂、ゴム弾性を有するポリ マー、多糖類等を1種または2種以上の混合物として用 いることができる。また、多糖類の様にリチウムと反応 する官能基を有する結着剤は、例えばメチル化するなど 折法による面間隔(d002)が3.354~3.36 50 してその官能基を失活させておくことが望ましい。その 添加量としては、1~50重量%が好ましく、特に2~ 30重量%が好ましい。

【0013】フィラーとしては、電池性能に悪影響を及 ぼさない材料であれば何でも良い。通常、ポリプロピレ ン、ポリエチレン等のオレフィン系ポリマー、アエロジ ル、ゼオライト、ガラス、炭素等が用いられる。フィラ -の添加量は0~30重量%が好ましい。

【0014】電極活物質の集電体としては、構成された 電池において悪影響を及ぼさない電子伝導体であれば何 でもよい。例えば、正極用集電体としては、アルミニウ 10 -の中に溶かし込んだものを用いることができる。さら ム、チタン、ステンレス鋼、ニッケル、焼成炭素、導電 性高分子、導電性ガラス等の他に、接着性、導電性、耐 酸化性向上の目的で、アルミニウムや銅等の表面をカー ボン、ニッケル、チタンや銀等で処理したものを用いる **ことができる。負極用集電体としては、銅、ステンレス** 鋼、ニッケル、アルミニウム、チタン、焼成炭素、導電 性高分子、導電性ガラス、AI-Cd合金等の他に、接 着性、導電性、耐酸化性向上の目的で、銅等の表面をカ ーボン、ニッケル、チタンや銀等で処理したものを用い 理することも可能である。これらの形状については、フ ォイル状の他、フィルム状、シート状、ネット状、パン チ又はエキスパンドされたもの、ラス体、多孔質体、発 泡体、繊維群の形成体等が用いられる。厚みは特に限定 はないが、 $1\sim500\mu$ mのものが用いられる。

【0015】正極活物質の主構成物質にLi,_, [Mn 1-x-v Mx LivO,]を用いる場合、他の正極活物質 を添加することも可能である。例えばMnO,, MoO ,, V, O,, Li, CoO,, Li, NiO, 等の金 属酸化物や、TiSz, MoSz, NbSe, 等の金属 カルコゲン化物、ポリアセン、ポリバラフェニレン、ポ リピロール、ポリアニリン、ジスルフィド等のグラファ イト層間化合物、及び導電性高分子等のアルカリ金属イ オンや、アニオンを吸放出可能な各種の物質を利用する ことができる。添加方法としては、単に混合する以外 に、電解重合、メッキ、焼結、メカノフュージョン、蒸 着等の技術を応用してコートすることが挙げられる。

【0016】また、電解質としては、例えば有機電解 液、高分子固体電解質、無機固体電解質、溶融塩等を用 いることができ、この中でも有機電解液を用いることが 40 好ましい。この有機電解液の有機溶媒として、プロピレ ンカーボネート、エチレンカーボネート、プチレンカー ボネート、ジエチルカーボネート、ジメチルカーボネー ト、メチルエチルカーボネート、アープチロラクトン等 のエステル類や、テトラヒドロフラン、2-メチルテト ラヒドロフラン等の置換テトラヒドロフラン、ジオキソ ラン、ジエチルエーテル、ジメトキシエタン、ジエトキ シエタン、メトキシエトキシエタン等のエーテル類、ジ メチルスルホキシド、スルホラン、メチルスルホラン、

ピロリドン、ジメチルフォルムアミド等が挙げられ、こ れらを単独又は混合溶媒として用いることができる。ま た、支持電解質塩としては、LiClO。、LiP F, LiBF, LiAsF, LiCF, SO, Lin(CF, SO,), Lin(C, F, SO,) ,、LiN(CF, SO,)(C, F, SO,), 等が 挙げられる。一方、高分子固体電解質としては、上記の ような支持電解質塩をボリエチレンオキシドやその架橋 体、ポリフォスファゼンやその架橋体等といったポリマ に、Li, N, Li I 等の無機固体電解質も使用可能で ある。つまり、リチウムイオン導伝性の非水電解質であ ればよい。

【0017】セパレータとしては、イオンの透過度が優 れ、機械的強度のある絶縁性薄膜を用いることができ る。耐有機溶剤性と疎水性からポリプロピレンやポリエ チレンといったオレフィン系のポリマー、ガラス繊維、 ポリフッ化ビニリデン、ポリテトラフルオロエチレン等 からつくられたシート、微孔膜、不織布、布が用いられ るととができる。これらの材料については表面を酸化処 20 る。セパレータの孔径は、一般に電池に用いられる範囲 のものであり、例えば0.01~10μmである。ま た、その厚みについても同様で、一般に電池に用いられ る範囲のものであり、例えば5~300μmである。 【0018】充放電特性、特にサイクル特性が向上する 理由として、必ずしも明確ではないが以下のように考察 される。一般的に、電池内部において、電池の充放電に 関与しない種々の不純物を含んでいることが多い。例え は、LiPF。を電解質に用いる場合、塩そのものが不 純物を持ち込んだり、電池内部や溶媒中に含まれる極微 量の水と反応することでHF(フッ酸)を生じることが 考えられる。リチウム吸蔵の際に炭素材料表面では、電 解液と炭素材料の間に炭酸リチウムのようなイオン伝導 性の高い被膜を形成するが、この被膜形成時あるいは形 成後にフッ酸の様な酸が存在すると、イオン伝導性の低 いハロゲン化リチウムを生じる。炭素材料と電解液の界 面に生じたハロゲン化リチウムは、リチウムの吸蔵放出 を妨げ、その結果負極の界面抵抗を増大させ、放電容量 が低減する原因の一つと考えられる。そこで、電池内部 に存在するフッ酸の量を低減することにより、この問題 が解決できるのではないかと考えた。つまり、このフッ 酸は電解質と水との反応によって生じるものと考えてい るが、正極活物質であるマンガン酸リチウムが触媒的に 分解を促進しているのではないかと考え、マンガンの一 部をリチウムとマンガン以外の元素で置換したスピネル 構造を有するマンガン酸リチウムを用いることにより、 分解等に及ぼす触媒的な活性を落とし、水と電解質との 分解反応によって生じるフッ酸を低減すると考た。さら に、この触媒的な活性度は、スピネル構造を有するLi Mn, O, の充電末物質である γ -MnO, が最も高 アセトニトリル、ギ酸メチル、酢酸メチル、N-メチル 50 く、マンガンの一部をリチウムとマンガン以外の元素で

置換することにより、このャーMn〇、の生成を抑制し 触媒活性が低減することができると考え、マンガンの一 部をリチウムとマンガン以外の元素で置換したスピネル 構造を有するマンガン酸リチウムを用いることで、負極 の炭素材料の界面抵抗増大が抑制され、サイクル特性が 向上したため、本発明に至った。また、このマンガンの 一部をリチウムとマンガン以外の元素で置換したスピネ ル構造を有するマンガン酸リチウムは、触媒活性が低減 しただけでなく、高温に於ける活物質自身の安定性も向 上し、高温に於けるサイクル特性も向上することが分か 10 った。

[0019]

【実施例】以下、本発明を実施例に基づき説明する。 【0020】(実施例1)酢酸リチウム二水和物と酢酸 マンガン(II)四水和物と酢酸カルシウム一水和物と をLiとMnとCaの比が1.10:1.95:0.0 5になるように混合し、これを酢酸に溶解した。熱を加 えながら撹拌し、完全に溶解した溶液から酢酸を蒸発さ せ、混合塩を得た。この混合塩を500℃で仮焼成し、 空気中850℃で焼成した。得られた焼成物を粉砕し、 このXRD測定を行った結果、スピネル構造を有するマ ンガン酸リチウムが得られていることが分かった。次に エネルギー分散型電子プローブマイクロアナリシス(E PMA) によりカルシウムの分散状態を観察したとと ろ、カルシウムはマンガン酸リチウムの全面に分布して いた。この粉末を粉末Aとする。

【0021】(実施例2)酢酸リチウム二水和物と酢酸 マンガン(II)四水和物と酢酸クロム(III)とを LiとMnとCrの比が1、10:1、95:0、05 になるように混合し、これを酢酸に溶解した。熱を加え 30 ながら撹拌し、完全に溶解した溶液から酢酸を蒸発さ せ、混合塩を得た。との混合塩を500℃で仮焼成し、 空気中850℃で焼成した。得られた焼成物を粉砕し、 このXRD測定を行った結果、スピネル構造を有するマ ンガン酸リチウムが得られていることが分かった。次に EPMAによりクロムの分散状態を観察したところ、ク ロムはマンガン酸リチウムの全面に分布していた。この 粉末を粉末Bとする。

【0022】 (実施例3) 酢酸リチウム二水和物と酢酸 マンガン([])四水和物と酢酸インジウム([]]) 水和物とをしiとMnとInの比が1.10:1.9 5:0.05になるように混合し、これを酢酸に溶解し た。熱を加えながら撹拌し、完全に溶解した溶液から酢 酸を蒸発させ、混合塩を得た。この混合塩を500℃で 仮焼成し、空気中850℃で焼成した。得られた焼成物 を粉砕し、このXRD測定を行った結果、スピネル構造 を有するマンガン酸リチウムが得られていることが分か った。次にEPMAによりインジウムの分散状態を観察 したところ、インジウムはマンガン酸リチウムの全面に 分布していた。この粉末を粉末Cとする。

【0023】(実施例4)酢酸リチウム二水和物と酢酸 マンガン(II)四水和物と酢酸テルビウム(III) 四水和物とをLiとMnとTbの比が1.10:1.9 5:0.05になるように混合し、これを酢酸に溶解し た。熱を加えながら撹拌し、完全に溶解した溶液から酢 酸を蒸発させ、混合塩を得た。この混合塩を500℃で 仮焼成し、空気中850℃で焼成した。得られた焼成物 を粉砕し、このXRD測定を行った結果、スピネル構造 を有するマンガン酸リチウムが得られていることが分か った。次にEPMAによりテルビウムの分散状態を観察 したところ、テルビウムはマンガン酸リチウムの全面に 分布していた。この粉末を粉末Dとする。

【0024】 (実施例5) 酢酸リチウム二水和物と酢酸 マンガン(【Ⅰ】)四水和物と酢酸マグネシウム(【Ⅰ】) 四水和物とをLiとMnとMgの比が1.10:1.9 5:0.05になるように混合し、これを酢酸に溶解し た。熱を加えながら撹拌し、完全に溶解した溶液から酢 酸を蒸発させ、混合塩を得た。この混合塩を500℃で 仮焼成し、空気中850°Cで焼成した。得られた焼成物 を粉砕し、このXRD測定を行った結果、スピネル構造 を有するマンガン酸リチウムが得られていることが分か った。次にEPMAによりクロム元素の分散状態を観察 したところ、マグネシウムはマンガン酸リチウムの全面 に分布していた。この粉末を粉末Eとする。

【0025】(実施例6)上記実施例1で得られた粉末 A、B, C, D, Eを正極活物質として用い、次のよう にして図1に示すコイン型非水電解質電池を試作した。 正極1は、粉末A~粉末Eとアセチレンブラック及びポ リテトラフルオロエチレン粉末とを重量比85:10: 5で混合し、トルエンを加えて十分混錬した。これをロ ーラープレスにより厚み0.8mmのシート状に成形し た。次にこれを直径16mmの円形に打ち抜き、減圧下 200℃で15時間乾燥し正極1を得た。正極1は正極 集電体6の付いた正極缶4に圧着して用いた。負極活物 質として、人造黒鉛(平均粒径6 μm、X線回折法によ る面間隔(d002)が3.37点で、C軸方向の結晶 の大きさ(Lc)が550Åとポリテトラフルオロエチ レン粉末とを重量比95:5で混合し、トルエンを加え て十分混練した。これをローラープレスにより厚み0. 1mmのシート状に成形した。次にこれを直径16mm の円形に打ち抜き、減圧下200℃で15時間乾燥して 負極2を得た。負極2は負極集電体7の付いた負極缶5 に圧着して用いた。エチレンカーボネートとジエチルカ ーボネートとの体積比1:1の混合溶剤にLiPF。を 1mol/1溶解した電解液を用い、セパレータ3には ポリプロピレン製微多孔膜を用いた。上記正極、負極、 電解液及びセパレータを用いて直径20mm、厚さ1. 6mmのコイン型リチウム電池を作製した。この粉末A ~Eを用いた電池をそれぞれ電池(A)~電池(E)と 50 する。

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【0026】(比較例)酢酸リチウム二水和物と酢酸マ * ンガン(II)四水和物とをLiとMnの比が1.1 0:2.00になるように混合し、これを酢酸に溶解した。熱を加えながら撹拌し、完全に溶解した溶液から酢酸を蒸発させ、混合塩を得た。この混合塩を500℃で仮焼成し、空気中850℃で焼成した。得られた焼成物を粉砕し、このXRD測定を行った結果、スピネル構造を有するマンガン酸リチウムが得られていることが分かった。この粉末を粉末Cとする。粉末Aの代わりに、粉末Fを用い、それ以外は実施例3と同様にして電池を作*10

*製した。得られた電池を比較電池(F)とする。 【0027】 これらの電池(A)~(E)、及び比較電池(F)を用いて充放電試験を行なった。充電終止電圧を4.2 V、放電終止電圧を3.0 Vとし、充放電電流を1mA、試験温度は20℃と40℃で定電流充放電を行った。得られた5サイクル目の放電容量の結果を表1に示した。また、サイクル寿命として放電容量が初期の60%に低下した時点のサイクル数を測定した。

[0028]

【表1】

包池	20℃での 放電容量 (mAh)	サイクル 対 命 (サイクル数)	40℃での 放電容量 (mAh)	サイクル 寿命 (サイクル数)
電池A	1 6	285	1 7	1 4 5
- 電池 B	1 6	2 6 5	17	140
電池C	16	3 0 0	1 7	1 6 0
電池D	1 6	265.	17	1 4 0
電池E	1 6	275	1 7	1 4 5
比較電池F	1 7	8 0	1.4	1 0

【0029】粉末A~粉末Eを用いた電池(A)~電池 30 (E) と比較電池 (F) を比較して初期の放電容量は変 わらなかったが、サイクル寿命が良くなることが分か る。また、40℃の高温においては、初期の放電容量は 増大し、サイクル寿命も向上することが分かる。つま り、本発明の正極活物質を用いることにより、電解質の 分解が少なく、これによってフッ酸の生成が抑制される ため、炭素表面に生成する表面被膜において、フッ酸の 存在下生成する抵抗の高いフッ化リチウムではなく、フ ッ素の関与しない比較的抵抗の低い炭酸リチウムや酸化 リチウムといったような成分の被膜が形成され、界面抵 40 抗増大が抑制されサイクル特性が向上することが考えら れる。また、高温の40℃でのサイクル特性も優れてい ることから、電解質の分解だけでなく、正極活物質自身 の電解液へのマンガンの溶出も抑えられ、活物質の容量 低下も抑制されることが考えられる上記実施例において は、正極活物質における主構成物質にLi[MniesC a..., Li., O.], Li [Mn.,, Cr.,, Li 0.10 O,], Li [Mn_{1.95} In_{0.05} Li_{0.10} O₄], Li [Mn_{1.9}, Tb_{9.0}, Li_{0.10}O,] Li [Mn_{1.9}, Mg.,,Li.,O,]を、負極材料として人造黒鉛を 50

) 用いたリチウム二次電池について挙げたが、同様の効果が他の置換元素及び負極材料についても確認された。なお、本発明は上記実施例に記載された活物質の出発原料、製造方法、正極、負極、電解質、セパレータ及び電池形状などに限定されるものではない。

[0030]

【発明の効果】本発明は上述の如く構成されているので、正極活物質の主構成物質であるスピネル構造を有するマンガン酸リチウムのマンガンの一部をリチウムとマンガン以外の元素で置換することにより、負極活物質である炭素材料界面での抵抗増大が少なく、さらに正極活物質自身の熱安定性が向上しサイクル特性が向上する。また、その材料が安全性に優れ、安価であることから、正極材料の優れた改質の方法であり、その結果得られる電池は、高容量、高エネルギー密度で、高温においても優れた充放電サイクル特性を示す。

【図面の簡単な説明】

【図1】本発明の実施例に係るコイン型非水電解質電池 の断面図である。

【符号の説明】

0 1 正極

12

負極

2 セパレータ 3

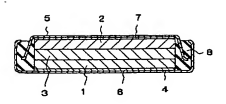
正極缶 4

***** 5 負極缶

> 6 正極集電体

負極集電体 7

(図1)



フロントページの続き

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CLAIMS <u>DETAILED DESCRIPTION TECHNICAL FIELD PRIOR ART EFFECT OF THE INVENTION TECHNICAL PROBLEM MEANS EXAMPLE DESCRIPTION OF DRAWINGS DRAWINGS</u>

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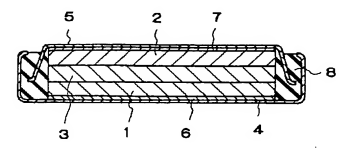
CLAIMS

[Claim(s)]

[Claim 1] It sets to the lithium secondary battery using a carbon material as a main constituent of a negative-electrode active material, and is Li1-z [Mn2-x-y Mx Liy O4] (M is elements other than Mn and Li, and x and y show the amount of substitution of manganese, and are 0 < x + y <= 1.) to the main constituent of a positive active material. Moreover, z shows the amount of lithiums usable in reversible, and is -1<=z<=1. Lithium secondary battery characterized by using.

[Claim 2] The main constituent of the aforementioned positive active material is Li1-z [Mn2-x-y CaxLiy O4] (x and y show the amount of substitution of manganese, and are $0 \le x + y \le 1$.). Moreover, z shows the amount of lithiums usable in reversible, and is -1<=z<=1. Lithium secondary battery according to claim 1 characterized by using. [Claim 3] The main constituent of the aforementioned positive active material is Li1-z [Mn2-x-y CrxLiy O4] (x and y show the amount of substitution of manganese, and are 0<x+y<=1.). Moreover, z shows the amount of lithiums usable in reversible, and is -1<=z<=1. Lithium secondary battery according to claim 1 characterized by using. [Claim 4] The main constituent of the aforementioned positive active material is Li1-z [Mn2-x-y InxLiy O4] (x and y show the amount of substitution of manganese, and are $0 \le x + y \le 1$.). Moreover, z shows the amount of lithiums usable in reversible, and is -1<=z<=1. Lithium secondary battery according to claim 1 characterized by using. [Claim 5] The main constituent of the aforementioned positive active material is Li1-z [Mn2-x-y TbxLiy O4] (x and y show the amount of substitution of manganese, and are 0 < x+y <= 1.). Moreover, z shows the amount of lithiums usable in reversible, and is -1<=z<=1. Lithium secondary battery according to claim 1 characterized by using. [Claim 6] The main constituent of the aforementioned positive active material is Li1-z [Mn2-x-y MgxLiy O4] (x and y show the amount of substitution of manganese, and are $0 \le x + y \le 1$.). Moreover, z shows the amount of lithiums usable in reversible, and is -1<=z<=1. Lithium secondary battery according to claim 1 characterized by using.

[Claim 7] The lithium secondary battery according to claim 1 characterized by the size (Lc) of the crystal of the spacing (d002) according [the carbon material which is the aforementioned negative-electrode active material] to an X-ray diffraction method of C shaft orientations by 3.354-3.369A being 200A or more.



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DETAILED DESCRIPTION

[Detailed Description of the Invention] [0001]

[The technical field to which invention belongs] this invention relates to a lithium secondary battery, and service capacity and power density are size and it relates to the high positive electrode for lithium secondary batteries and high negative electrode of the safety which was especially excellent in the cycle property in an elevated temperature.

[0002]

[Description of the Prior Art] as the positive active material of the lithium secondary battery characterized by 4V now -- LiCoO2 and LiNiO2 etc. -- alpha-NaFeO 2 The ** lithium oxide which has structure, or LiMn 2O4 etc. -the ** lithium oxide which has Spinel structure is used conventionally LiMn 204 which has Spinel structure especially It is the positive active material excellent in a low cost and safety. On the other hand, as a negativeelectrode active material, although the lithium metal and the lithium alloy had been used conventionally, that the short circuit and cycle life of positive/negative two poles by deposit (dendrite) of an arborescence lithium are short, and a part for the degradation should be compensated for the reason, the lithium of the 3 time equivalent of cell capacity is required for these cells, and they had a fault of a low in an energy density. In order to solve these troubles recently, the research which uses a carbon material for a negative electrode is active. When using the graphite to which this kind of negative electrode, especially graphitization progressed (for example, if a manganic acid lithium is used for a positive electrode, when a cell voltage will become a flat and will use for the pocket device of cell use), it is predominant in respect of capacity. However, it is LiMn 2O4 to a positive electrode about negative-electrode graphite. If it uses and a charge-and-discharge cycle is performed, if the dope voltage of the graphite at the time of charge piles up a cycle, it will become the 0V neighborhood, and will become competitive reaction with a deposit of a lithium. Therefore, cycle degradation was set to one of the large factors. Moreover, since this manganic acid lithium is not excellent in the cycle property of a positive active material own [the], replacing some manganese of the manganic acid lithium of Spinel structure by elements other than manganese in JP,4-233161,A, JP,5-21067,A, or JP,6-187993,A is reported. However, since manganese was eluted in the electrolytic solution when it was left at the elevated temperature also in such material, the charge-and-discharge capacity of a positive electrode decreased, and it turns out that cycle degradation is carried out. [0003]

[Problem(s) to be Solved by the Invention] As mentioned above, when a carbon material is used as a negative electrode, there is a problem of the increase in a carbonaceous interfacial resistance with combination with a manganic acid lithium. Moreover, there is also a problem that charge-and-discharge capacity falls by the charge and discharge in the elevated temperature of a positive active material itself. this invention is Li1-z [Mn2-x-y Mx Liy O4] (x and y show the amount of substitution of manganese, and are 0 < x + y < 1.) to the main constituent [in / a positive active material / since this trouble is solved]. Moreover, z shows the amount of lithiums usable in reversible, and is -1<=z<=1. By using, also in combination with a carbon material, or an elevated temperature, it is high capacity and high-energy density, and aims at offering the lithium secondary battery which was excellent in the charge-and-discharge cycle property with little irreversible capacity.

[Means for Solving the Problem] Although the occlusion of the lithium to a carbon material and discharge (an intercalation, day intercalation) were the reactions which mainly occur when carbon was considered as a negativeelectrode active material, it turns out that the coat state produced between the electrolytic solution and a carbon front face is involving as one of the factors which govern the reaction. For example, it is precise, the high coat of ion conductivity is excellent also in the cell property, and it is known that an ion conductivity conversely thick low coat has bad rate property and cycle property so that it may be represented with the case where a lithium metal is made into a negative-electrode active material. In this case, the former is coats, such as a lithium carbonate and a lithium oxide, and it is reported that the latters are coats, such as lithium fluoride. The coat which the same thing as

this produces on a carbon front face is also considered. That is, formation of the low coat of ionic conductivity, such as lithium fluoride, is raised on the surface of a carbon material as one of the factors which increases the interfacial resistance of a carbon material. In process in which the coat of this lithium fluoride is formed, it is possible that the fluoric acid which an electrolyte decomposes with the water carried in from positive-electrode material etc., and is generated is involving. This invention persons found out that the fluoric acid generated inside a cell could be suppressed in the manganic acid lithium which has Spinel structure by replacing some manganese by elements other than manganese, such as a lithium, calcium and chromium, an indium, a terbium, and magnesium, as a result of examining many things, in order to solve this trouble. Moreover, it turns out that this positive-electrode material also suppresses elution of the manganese to the electrolytic solution by elevated-temperature preservation. [0005] Li1-z which is the main constituent of a positive active material In [Mn2-x-y Mx Liy O4], M is elements other than Mn and Li, and its element which can be replaced by Mn is desirable. For example, Be, B, C, Si, P, Sc, Cu, Zn, Ga, germanium, As, Se, Sr, Mo, Pd, Ag, Cd, In, Sn, Sb, Te, Ba, Ta, W.Pb, Bi, Co, Fe, Cr, nickel, Ti, Zr, Nb, Y, aluminum, Na, K, Mg, calcium, Cs, La, Ce, Nd, Sm, Eu, Tb, etc. are mentioned. Also in these, since the effect of Cr, calcium, In, Tb, and Mg was remarkable, they are the most desirable. Here, x and y which show the amount of substitution of Mn are $0 \le x + y \le 1$. however, the thing replaced in $0 \le x + y \le 0.2$ in order that z which shows the amount of lithiums usable in reversible may decrease, if amount x+y of substitution of Mn is increased -good -- better Although z which shows the amount of lithiums usable in reversible changes also with amounts of substitution of Mn, it is $-1 \le z \le 1$.

[0006] Li1-z which is the main constituent of a positive active material The method of adding the element beforehand replaced by the baking raw material in [Mn2-x-y Mx Liy O4], when a lithium and a different-species element replace a part of Mn, and LiMn 2O4 Although the way the ion exchange etc. replaces a different-species element etc. is mentioned after calcinating, it is not limited to these.

[0007] The spacing (d002) especially by the X-ray diffraction method is 3.354 to 3.369 A, and since high capacity is obtained, the carbon particle whose size (Lc) of the crystal of C shaft orientations is 200A or more is [that the carbon material used for a negative-electrode active material should just be a carbon material which emits / occlusion and / a lithium] desirable.

[0008] As for the positive electrode used for this invention, and a negative-electrode active material, it is desirable that it is 100 micrometers or less of average grain size. When acquiring a predetermined configuration, a grinder and a classifier are used in order to obtain fine particles. For example, a mortar, a ball mill, a sand mill, a vibration ball mill, a planet ball mill, a jet mill, a counter jet mill, a revolution air current type jet mill, a thermal plasma, a screen, etc. are used. At the time of trituration, wet grinding which made organic solvents, such as water or a hexane, live together can also be used. As the classification method, there is especially no limitation and a screen, a pneumatic elutriation machine, etc. are used if needed in dry type and wet.

[0009] As a negative-electrode material which can be combined and can be used for this invention, the organic compound containing lithiums, such as a lithium metal, a lithium alloy, etc. and a chalcogen compound, a methyl lithium, etc. is mentioned. Moreover, it is also possible by using together the organic compound containing a lithium metal, a lithium alloy, and a lithium to insert a lithium in the carbon material used for this invention beforehand.

[0010] the case where the positive electrode of this invention and a negative-electrode active material are used -- an electrode -- an electric conduction agent, a binder, a filler, etc. can be added as a mixture If it is the electronic-conduction nature material which does not have a bad influence on a cell performance as an electric conduction agent, it is good anything. Usually, conductive material, such as natural graphites (a flaky graphite, a scale-like graphite, earthy graphite, etc.), an artificial graphite, carbon black, acetylene black, KETCHIEN black, a carbon whisker, carbon fiber metallurgy group powder (copper, nickel, aluminum, silver, gold, etc.), a metal fiber, and a conductive ceramic material, can be included as one sort or those mixture. In these, combined use of acetylene black and KETCHIEN black is desirable. The addition has 1 - 50 desirable % of the weight, and its 2 - 30 % of the weight is especially desirable.

[0011] When using the positive electrode of this invention, and a negative-electrode active material, the thing of the fine particles for which it is things other than the main active material, and a surface-layer portion is embellished at least is also possible. For example, applying technology, such as plating, sintering, a mechano fusion, and vacuum evaporationo, and carrying out the coat of the ion conductivity good matter, such as good matter of electronic-conduction nature, such as gold, silver, carbon, nickel, and copper, and a lithium carbonate, boron glass, a solid electrolyte, is mentioned.

[0012] As a binder, thermoplastics, such as a tetrafluoroethylene, a polyvinylidene fluoride, polyethylene, polypropylene, an ethylene-propylene-diene terpolymer (EPDM), sulfonation EPDM, styrene butadiene rubber (SBR), a fluororubber, and a carboxymethyl cellulose, the polymer which has rubber elasticity, polysaccharide, etc. can usually be used as one sort or two sorts or more of mixture. Moreover, as for a lithium and the binder which has the functional group which reacts, it is desirable like polysaccharide to methylate, for example and to make the

functional group deactivate. As the addition, 1 - 50 % of the weight is desirable, and 2 - 30 % of the weight is especially desirable.

[0013] If it is the material which does not have a bad influence on a cell performance as a filler, it is good anything. Usually, olefin system polymer, such as polypropylene and polyethylene, Aerosil, a zeolite, glass, carbon, etc. are used. The addition of a filler has 0 - 30 desirable % of the weight.

[0014] If it is the electronic-conduction object which does not do a bad influence in the constituted cell as a charge collector of an electrode active material, it is good anything. For example, as a charge collector for positive electrodes, besides aluminum, titanium, stainless steel, nickel, a baked carbon, a conductive polymer, electrically conductive glass, etc., it is the purpose of an adhesive property, conductivity, and oxidation-resistant improvement, and what processed front faces, such as aluminum and copper, with carbon, nickel, titanium, silver, etc. can be used. As a charge collector for negative electrodes, besides copper, stainless steel, nickel, aluminum, titanium, a baked carbon, a conductive polymer, electrically conductive glass, an aluminum-Cd alloy, etc., it is the purpose of an adhesive property, conductivity, and oxidation-resistant improvement, and what processed front faces, such as copper, with carbon, nickel, titanium, silver, etc. can be used. About such material, it is also possible to oxidize a front face. About these configurations, the shape of the shape of others and a film and a sheet, the letter of a network, punch or the thing by which expanded one was carried out, a lath object, a porosity object, a foam, the organizer of a fiber group, etc. are used. [shape / of foil] Although especially limitation is thin, a 1-500-micrometer thing is used.

[0015] It is Li1-z to the main constituent of a positive active material. When using [Mn2-x-y Mx Liy O4], it is also possible to add other positive active materials. for example, MnO2, MoO3, V2 O5, Lix CoO2, and Lix NiO2 etc. -a metallic oxide, and TiS2, MoS2 and NbSe3 etc. -- various kinds of matter in which absorption/emission is possible can be used for alkali-metal ion, such as intercalated graphite, such as a metal chalcogen ghost, the poly acene, poly para-phenylene, polypyrrole, the poly aniline, and disulfide, and a conductive polymer, and an anion Applying and carrying out the coat of the technology, such as electrolytic polymerization, plating, sintering, a mechano fusion, and vacuum evaporationo, as the addition method besides only mixing is mentioned. [0016] Moreover, it is desirable to be able to use the organic electrolytic solution, a solid polymer electrolyte, an inorganic solid electrolyte, fused salt, etc., for example, and to use the organic electrolytic solution also in this as an electrolyte. As an organic solvent of this organic electrolytic solution, propylene carbonate, ethylene carbonate, Butylene carbonate, diethyl carbonate, dimethyl carbonate, Ester, such as methylethyl carbonate and gammabutyrolactone Substitution tetrahydrofurans, such as a tetrahydrofuran and 2-methyl tetrahydrofuran, A dioxolane, diethylether, dimethoxyethane, diethoxy ethane, Ether, such as methoxyethoxy ethane, dimethyl sulfoxide, a sulfolane, a methyl sulfolane, an acetonitrile, methyl formate, methyl acetate, N-methyl pyrrolidone, a dimethyl formamide, etc. are mentioned, and these can be used as independent or a mixed solvent. moreover -- as a supporting-electrolyte salt -- LiClO4, LiPF6, LiBF4, LiAsF6, LiCF3 SO3, LiN (CF3 SO2)2, LiN (C2 F5 SO2)2, and LiN (CF3 SO2) (C4 F9 SO2)2 etc. -- it is mentioned On the other hand, what melted the above supportingelectrolyte salts as a solid polymer electrolyte into polymer, such as a polyethylene oxide, the bridge formation object and poly force FAZEN, and its bridge formation object, can be used. Furthermore, inorganic solid electrolytes, such as Li3 N and LiI, are also usable. That is, what is necessary is just nonaqueous electrolyte of lithium ion conductivity.

[0017] As separator, the transmittance of ion is excellent and an insulating thin film with a mechanical strength can be used. The sheet built with the polymer of olefin systems, such as organic-solvent-proof nature, hydrophobic shell polypropylene, and polyethylene, the glass fiber, the polyvinylidene fluoride, the polytetrafluoroethylene, etc., a micropore film, a nonwoven fabric, and cloth are used. The aperture of separator is a thing of the range generally used for a cell, for example, is 0.01-10 micrometers. Moreover, the same is said of the thickness, and it is the thing of the range generally used for a cell, for example, is 5-300 micrometers.

[0018] As a reason a charge-and-discharge property, especially whose cycle property improve, although it is not necessarily clear, it is considered as follows. Generally, in the interior of a cell, the various impurities which do not participate in the charge and discharge of a cell are included in many cases. For example, LiPF6 When using for an electrolyte, the salt itself can carry in an impurity or it is possible to produce HF (fluoric acid) reacting with the water of the ultralow volume contained in the interior of a cell, or a solvent. Although an ion conductivity high coat like a lithium carbonate is formed between the electrolytic solution and a carbon material on a carbon-material front face in the case of lithium occlusion, if an acid like fluoric acid exists after the time of this coat formation, or formation, an ion conductivity low lithium halide will be produced. The lithium halide produced in the interface of a carbon material and the electrolytic solution bars occlusion discharge of a lithium, as a result, increases the interfacial resistance of a negative electrode, and is considered to be one of the causes which service capacity reduces. Then, I thought that this problem would be solvable by reducing the amount of the fluoric acid which exists in the interior of a cell. That is, by the manganic acid lithium which is a positive active material thinking that decomposition will be promoted in catalyst, and using the manganic acid lithium which has the Spinel structure

which replaced some manganese by elements other than a lithium and manganese, although it thinks that it is generated by the reaction of an electrolyte and water, this fluoric acid is ****, when the fluoric acid which drops the catalyst-activity exerted on decomposition etc. and is produced by the decomposition reaction of water and an electrolyte is reduced. Furthermore, this catalyst-activity is LiMn 2O4 which has Spinel structure. Gamma-MnO 2 which is the matter after charge By being the highest and replacing some manganese by elements other than a lithium and manganese This gamma-MnO 2 By using the manganic acid lithium which has the Spinel structure which suppressed generation, thought that catalytic activity could be reduced and replaced some manganese by elements other than a lithium and manganese Since interfacial-resistance increase of the carbon material of a negative electrode was suppressed and the cycle property improved, it resulted in this invention. Moreover, it turns out that its stability of an active material own [hot] improves catalytic activity not only reduced the manganic acid lithium which has the Spinel structure which replaced a part of this manganese by elements other than a lithium and manganese, but, and a hot cycle property also improves.

[Example] Hereafter, this invention is explained based on an example.

[0020] (Example 1) The acetic-acid lithium dihydrate, manganese acetate (II)4 hydrate, and the calcium-acetate monohydrate were mixed so that Li and the ratio of Mn and calcium might be set to 1.10:1.95:0.05, and this was dissolved in the acetic acid. It agitated applying heat, the acetic acid was evaporated from the solution which dissolved completely, and the mixed salt was obtained. Temporary baking of this mixed salt was carried out at 500 degrees C, and it calcinated at 850 degrees C among air. As a result of grinding the obtained baking object and performing this XRD measurement, it turns out that the manganic acid lithium which has Spinel structure is obtained. Next, when the distributed state of calcium was observed by energy-dispersion type electron probe microanalysis (EPMA), calcium was distributed all over the manganic acid lithium. Let this powder be Powder A. [0021] (Example 2) The acetic-acid lithium dihydrate, manganese acetate (II)4 hydrate, and the chromium acetate (III) were mixed so that the ratio of Li, and Mn and Cr might be set to 1.10:1.95:0.05, and this was dissolved in the acetic acid. It agitated applying heat, the acetic acid was evaporated from the solution which dissolved completely, and the mixed salt was obtained. Temporary baking of this mixed salt was carried out at 500 degrees C, and it calcinated at 850 degrees C among air. As a result of grinding the obtained baking object and performing this XRD measurement, it turns out that the manganic acid lithium which has Spinel structure is obtained. Next, when the distributed state of chromium was observed by EPMA, chromium was distributed all over the manganic acid lithium. Let this powder be Powder B.

[0022] (Example 3) The acetic-acid lithium dihydrate, manganese acetate (II)4 hydrate, and the acetic-acid indium (III) hydrate were mixed so that the ratio of Li, and Mn and In might be set to 1.10:1.95:0.05, and this was dissolved in the acetic acid. It agitated applying heat, the acetic acid was evaporated from the solution which dissolved completely, and the mixed salt was obtained. Temporary baking of this mixed salt was carried out at 500 degrees C, and it calcinated at 850 degrees C among air. As a result of grinding the obtained baking object and performing this XRD measurement, it turns out that the manganic acid lithium which has Spinel structure is obtained. Next, when the distributed state of an indium was observed by EPMA, the indium was distributed all over the manganic acid lithium. Let this powder be Powder C.

[0023] (Example 4) The acetic-acid lithium dihydrate, manganese acetate (II)4 hydrate, and acetic-acid terbium (III) 4 hydrate were mixed so that Li and the ratio of Mn and Tb might be set to 1.10:1.95:0.05, and this was dissolved in the acetic acid. It agitated applying heat, the acetic acid was evaporated from the solution which dissolved completely, and the mixed salt was obtained. Temporary baking of this mixed salt was carried out at 500 degrees C, and it calcinated at 850 degrees C among air. As a result of grinding the obtained baking object and performing this XRD measurement, it turns out that the manganic acid lithium which has Spinel structure is obtained. Next, when the distributed state of a terbium was observed by EPMA, the terbium was distributed all over the manganic acid lithium. Let this powder be Powder D.

[0024] (Example 5) The acetic-acid lithium dihydrate, manganese acetate (II)4 hydrate, and magnesium-acetate (II)4 hydrate were mixed so that Li and the ratio of Mn and Mg might be set to 1.10:1.95:0.05, and this was dissolved in the acetic acid. It agitated applying heat, the acetic acid was evaporated from the solution which dissolved completely, and the mixed salt was obtained. Temporary baking of this mixed salt was carried out at 500 degrees C, and it calcinated at 850 degrees C among air. As a result of grinding the obtained baking object and performing this XRD measurement, it turns out that the manganic acid lithium which has Spinel structure is obtained. Next, when the distributed state of a chromium element was observed by EPMA, magnesium was distributed all over the manganic acid lithium. Let this powder be Powder E.

[0025] (Example 6) The coin type nonaqueous electrolyte cell shown in <u>drawing 1</u> as follows was made as an experiment, using the powder A, B, C, D, and E obtained in the above-mentioned example 1 as a positive active material. The positive electrode 1 mixed Powder A - Powder E, acetylene black, and polytetrafluoroethylene powder by the weight ratio 85:10:5, added toluene, and kneaded it enough. This was fabricated with a thickness of

0.8mm in the shape of a sheet with the roller press. next, this -- the diameter of 16mm -- it pierced circularly, and dried at 200 degrees C under reduced pressure for 15 hours, and the positive electrode 1 was obtained The positive electrode 1 was stuck by pressure and used for the positive-electrode can 4 to which the positive-electrode charge collector 6 was attached. As a negative-electrode active material, it is an artificial graphite (the size (Lc) of the crystal of the spacing (d002) by 6 micrometers of mean particle diameters and the X-ray diffraction method of C shaft orientations by 3.37A mixed 550A and polytetrafluoroethylene powder by the weight ratio 95:5, added toluene, and kneaded enough.). This was fabricated with a thickness of 0.1mm in the shape of a sheet with the roller press. next, this -- the diameter of 16mm -- it pierced circularly, it dried at 200 degrees C under reduced pressure for 15 hours, and the negative electrode 2 was obtained The negative electrode 2 was stuck by pressure and used for the negative-electrode can 5 to which the negative-electrode charge collector 7 was attached. It is LiPF6 to the partially aromatic solvent of the volume ratio 1:1 of ethylene carbonate and diethyl carbonate. The fine porous membrane made from polypropylene was used for separator 3 using the electrolytic solution which carried out 1 mol/l dissolution. The coin type lithium cell with a diameter [of 20mm] and a thickness of 1.6mm was produced using the above-mentioned positive electrode, a negative electrode, the electrolytic solution, and separator. Let the cells using this powder A-E be a cell (A) - a cell (E), respectively.

[0026] (Example of comparison) The acetic-acid lithium dihydrate and manganese acetate (II)4 hydrate were mixed so that the ratio of Li and Mn might be set to 1.10:2.00, and this was dissolved in the acetic acid. It agitated applying heat, the acetic acid was evaporated from the solution which dissolved completely, and the mixed salt was obtained. Temporary baking of this mixed salt was carried out at 500 degrees C, and it calcinated at 850 degrees C among air. As a result of grinding the obtained baking object and performing this XRD measurement, it turns out that the manganic acid lithium which has Spinel structure is obtained. Let this powder be Powder C. Instead of Powder A, Powder F was used and the cell was produced like the example 3 except it. Let the obtained cell be a comparison cell (F).

[0027] These cells (A) The charge and discharge test was performed using - (E) and the comparison cell (F). Charge final voltage was set to 4.2V, the discharge final voltage was set to 3.0V, and 1mA and the test temperature performed constant-current charge and discharge for the charge and discharge current at 20 degrees C and 40 degrees C. The result of the service capacity of obtained 5 cycle eye was shown in Table 1. Moreover, the number of cycles at the time of service capacity falling to 60% of the first stage as a cycle life was measured. [0028]

Ta	ble	: 1]	Ĺ

Table 1		· · · · · · · · · · · · · · · · · · ·		~
瑶 池	20℃での 放電容量 (mAh)	サイクル 対 命 (サイクル数)	40℃での 放電容量 (mAh)	サイクル寿命 (サイクル数)
電池A	16	285	1 7	1 4 5
電池B	1 6	2 6 5	1 7	1 4 0
電池C	. 16	300	17	1 6 0
電池D	1 6	265.	1 7	1 4 0
電池E	1 6	275	1 7	1 4 5
比較電池F	1 7	8 0	1 4	1 0

[0029] Although the cell (A) - cell (E), and comparison cell (F) using Powder A - Powder E were compared and early service capacity did not change, a cycle life is good and a bird clapper is known. Moreover, in the elevated temperature of 40 degrees C, it turns out that early service capacity increases and a cycle life also improves. That is, there is little electrolytic disassembly by using the positive active material of this invention, and since generation of fluoric acid is suppressed by this, in the surface lining generated on a carbon front face, it is possible [it] that the coat of the component like not the high lithium fluoride of the resistance generated under existence of fluoric acid but the lithium carbonate with comparatively low resistance in which a fluorine does not participate and a lithium

oxide is formed, interfacial-resistance increase is suppressed, and a cycle property improves. moreover, since the 40-degree C hot cycle property is also excellent, not only in electrolytic disassembly In the above-mentioned example which can consider that elution of the manganese to the own electrolytic solution of a positive active material is also suppressed, and capacity deterioration of an active material is also suppressed To the main constituent in a positive active material, Li [Mn1.95calcium0.05Li 0.10O4], Li [Mn1.95Cr0.05Li 0.10O4] and Li [Mn1.95In0.05Li 0.10O4], Li [Mn1.95Tb0.05Li 0.10O4] Although Li [Mn1.95Mg0.05Li 0.10O4] was mentioned about the lithium secondary battery using the artificial graphite as a negative-electrode material, the same effect was checked about other substitution elements and negative-electrode material. In addition, this invention is not limited to the start raw material, the manufacture method, a positive electrode, a negative electrode, an electrolyte, separator, a cell configuration, etc. of the active material indicated by the above-mentioned example.

[Effect of the Invention] Since this invention is constituted like ****, by replacing some manganese of the manganic acid lithium which has the Spinel structure which is the main constituent of a positive active material by elements other than a lithium and manganese, it has little resistance increase by the carbon-material interface which is a negative-electrode active material, its own thermal stability of a positive active material improves further, and its cycle property improves. Moreover, it is the method of reforming which the material was excellent in safety, and was excellent in positive-electrode material since it was cheap, and the cells obtained as a result are high capacity and high-energy density, and show the charge-and-discharge cycle property of having excelled also in the elevated temperature.

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TECHNICAL FIELD

[The technical field to which invention belongs] this invention relates to a lithium secondary battery, and service capacity and power density are size and it relates to the high positive electrode for lithium secondary batteries and high negative electrode of the safety which was especially excellent in the cycle property in an elevated temperature.

[0002]

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PRIOR ART

[Description of the Prior Art] as the positive active material of the lithium secondary battery characterized by 4V now -- LiCoO2 and LiNiO2 etc. -- alpha-NaFeO 2 The ** lithium oxide which has structure, or LiMn 2O4 etc. -the ** lithium oxide which has Spinel structure is used conventionally LiMn 2O4 which has Spinel structure especially It is the positive active material excellent in a low cost and safety. On the other hand, as a negativeelectrode active material, although the lithium metal and the lithium alloy had been used conventionally, that the short circuit and cycle life of positive/negative two poles by deposit (dendrite) of an arborescence lithium are short, and a part for the degradation should be compensated for the reason, the lithium of the 3 time equivalent of cell capacity is required for these cells, and they had the fault that an energy density was low. In order to solve these troubles recently, the research which uses a carbon material for a negative electrode is active. When using the graphite to which this kind of negative electrode, especially graphitization progressed (for example, if a manganic acid lithium is used for a positive electrode, when a cell voltage will become a flat and will use for the pocket device of cell use), it is predominant in respect of capacity. However, it is LiMn 204 to a positive electrode about negative-electrode graphite. If it uses and a charge-and-discharge cycle is performed, if the dope voltage of the graphite at the time of charge piles up a cycle, it will become the 0V neighborhood, and will become competitive reaction with a deposit of a lithium. Therefore, cycle degradation was set to one of the large factors. Moreover, since this manganic acid lithium is not excellent in the cycle property of a positive active material own [the], replacing some manganese of the manganic acid lithium of Spinel structure by elements other than manganese in JP,4-233161,A, JP,5-21067,A, or JP,6-187993,A is reported. However, since manganese was eluted in the electrolytic solution when it was left at the elevated temperature also in such material, the charge-and-discharge capacity of a positive electrode decreased, and it turns out that cycle degradation is carried out.

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EFFECT OF THE INVENTION

[Effect of the Invention] Since this invention is constituted like ****, by replacing some manganese of the manganic acid lithium which has the Spinel structure which is the main constituent of a positive active material by elements other than a lithium and manganese, it has little resistance increase by the carbon-material interface which is a negative-electrode active material, its own thermal stability of a positive active material improves further, and its cycle property improves. Moreover, it is the method of reforming which the material was excellent in safety, and was excellent in positive-electrode material since it was cheap, and the cells obtained as a result are high capacity and high-energy density, and show the charge-and-discharge cycle property of having excelled also in the elevated temperature.

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TECHNICAL PROBLEM

[Problem(s) to be Solved by the Invention] As mentioned above, when a carbon material is used as a negative electrode, there is a problem of the increase in a carbonaceous interfacial resistance with combination with a manganic acid lithium. Moreover, there is also a problem that charge-and-discharge capacity falls by the charge and discharge in the elevated temperature of a positive active material itself. this invention is Li1-z [Mn2-x-y Mx Liy O4] (x and y show the amount of substitution of manganese, and are 0<x+y<=1.) to the main constituent [in / a positive active material / since this trouble is solved]. Moreover, z shows the amount of lithiums usable in reversible, and is -1<=z<=1. By using, also in combination with a carbon material, or an elevated temperature, it is high capacity and high-energy density, and aims at offering the lithium secondary battery which was excellent in the charge-and-discharge cycle property with little irreversible capacity.

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MEANS

[Means for Solving the Problem] Although the occlusion of the lithium to a carbon material and discharge (an intercalation, day intercalation) were the reactions which mainly occur when carbon was considered as a negativeelectrode active material, it turns out that the coat state produced between the electrolytic solution and a carbon front face is involving as one of the factors which govern the reaction. For example, it is precise, the high coat of ion conductivity is excellent also in the cell property, and it is known that an ion conductivity conversely thick low coat has bad rate property and cycle property so that it may be represented with the case where a lithium metal is made into a negative-electrode active material. In this case, the former is coats, such as a lithium carbonate and a lithium oxide, and it is reported that the latters are coats, such as lithium fluoride. The coat which the same thing as this produces on a carbon front face is also considered. That is, formation of the low coat of ionic conductivity, such as lithium fluoride, is raised on the surface of a carbon material as one of the factors which increases the interfacial resistance of a carbon material. In process in which the coat of this lithium fluoride is formed, it is possible that the fluoric acid which an electrolyte decomposes with the water carried in from positive-electrode material etc., and is generated is involving. This invention persons found out that the fluoric acid generated inside a cell could be suppressed in the manganic acid lithium which has Spinel structure by replacing some manganese by elements other than manganese, such as a lithium, calcium and chromium, an indium, a terbium, and magnesium, as a result of examining many things, in order to solve this trouble. Moreover, it turns out that this positive-electrode material also suppresses elution of the manganese to the electrolytic solution by elevated-temperature preservation. [0005] Li1-z which is the main constituent of a positive active material In [Mn2-x-y Mx Liy O4], M is elements other than Mn and Li, and its element which can be replaced by Mn is desirable. For example, Be, B, C, Si, P, Sc, Cu, Zn, Ga, germanium, As, Se, Sr, Mo, Pd, Ag, Cd, In, Sn, Sb, Te, Ba, Ta, W.Pb, Bi, Co, Fe, Cr, nickel, Ti, Zr, Nb, Y, aluminum, Na, K, Mg, calcium, Cs, La, Ce, Nd, Sm, Eu, Tb, etc. are mentioned. Also in these, since the effect of Cr, calcium, In, Tb, and Mg was remarkable, they are the most desirable. Here, x and y which show the amount of substitution of Mn are 0 < x+y <= 1. however, the thing replaced in 0 < x+y <= 0.2 in order that z which shows the amount of lithiums usable in reversible may decrease, if amount x+y of substitution of Mn is increased -good -- better Although z which shows the amount of lithiums usable in reversible changes also with amounts of substitution of Mn, it is $-1 \le z \le 1$.

[0006] Li1-z which is the main constituent of a positive active material The method of adding the element beforehand replaced by the baking raw material in [Mn2-x-y Mx Liy O4], when a lithium and a different-species element replace a part of Mn, and LiMn 2O4 Although the way the ion exchange etc. replaces a different-species element etc. is mentioned after calcinating, it is not limited to these.

[0007] The spacing (d002) especially by the X-ray diffraction method is 3.354 to 3.369 A, and since high capacity is obtained, the carbon particle whose size (Lc) of the crystal of C shaft orientations is 200A or more is [that the carbon material used for a negative-electrode active material should just be a carbon material which emits / occlusion and / a lithium] desirable.

[0008] As for the positive electrode used for this invention, and a negative-electrode active material, it is desirable that it is 100 micrometers or less of average grain size. When acquiring a predetermined configuration, a grinder and a classifier are used in order to obtain fine particles. For example, a mortar, a ball mill, a sand mill, a vibration ball mill, a planet ball mill, a jet mill, a counter jet mill, a revolution air current type jet mill, a thermal plasma, a screen, etc. are used. At the time of trituration, wet grinding which made organic solvents, such as water or a hexane, live together can also be used. As the classification method, there is especially no limitation and a screen, a pneumatic elutriation machine, etc. are used if needed in dry type and wet.

[0009] As a negative-electrode material which can be combined and can be used for this invention, the organic compound containing lithiums, such as a lithium metal, a lithium alloy, etc. and a chalcogen compound, a methyl lithium, etc. is mentioned. Moreover, it is also possible by using together the organic compound containing a lithium metal, a lithium alloy, and a lithium to insert a lithium in the carbon material used for this invention beforehand.

[0010] the case where the positive electrode of this invention and a negative-electrode active material are used -- an electrode -- an electric conduction agent, a binder, a filler, etc. can be added as a mixture If it is the electronic-conduction nature material which does not have a bad influence on a cell performance as an electric conduction agent, it is good anything. Usually, conductive material, such as natural graphites (a flaky graphite, a scale-like graphite, earthy graphite, etc.), an artificial graphite, carbon black, acetylene black, KETCHIEN black, a carbon whisker, carbon fiber metallurgy group powder (copper, nickel, aluminum, silver, gold, etc.), a metal fiber, and a conductive ceramic material, can be included as one sort or those mixture. In these, combined use of acetylene black and KETCHIEN black is desirable. The addition has 1 - 50 desirable % of the weight, and its 2 - 30 % of the weight is especially desirable.

[0011] When using the positive electrode of this invention, and a negative-electrode active material, the thing of the fine particles for which it is things other than the main active material, and a surface-layer portion is embellished at least is also possible. For example, applying technology, such as plating, sintering, a mechano fusion, and vacuum evaporationo, and carrying out the coat of the ion conductivity good matter, such as good matter of electronic-conduction nature, such as gold, silver, carbon, nickel, and copper, and a lithium carbonate, boron glass, a solid electrolyte, is mentioned.

[0012] As a binder, thermoplastics, such as a tetrafluoroethylene, a polyvinylidene fluoride, polyethylene, polypropylene, an ethylene-propylene-diene terpolymer (EPDM), sulfonation EPDM, styrene butadiene rubber (SBR), a fluororubber, and a carboxymethyl cellulose, the polymer which has rubber elasticity, polysaccharide, etc. can usually be used as one sort or two sorts or more of mixture. Moreover, as for a lithium and the binder which has the functional group which reacts, it is desirable like polysaccharide to methylate, for example and to make the functional group deactivate. As the addition, 1 - 50 % of the weight is desirable, and 2 - 30 % of the weight is especially desirable.

[0013] If it is the material which does not have a bad influence on a cell performance as a filler, it is good anything. Usually, olefin system polymer, such as polypropylene and polyethylene, Aerosil, a zeolite, glass, carbon, etc. are used. The addition of a filler has 0 - 30 desirable % of the weight.

[0014] If it is the electronic-conduction object which does not do a bad influence in the constituted cell as a charge collector of an electrode active material, it is good anything. For example, as a charge collector for positive electrodes, besides aluminum, titanium, stainless steel, nickel, a baked carbon, a conductive polymer, electrically conductive glass, etc., it is the purpose of an adhesive property, conductivity, and oxidation-resistant improvement, and what processed front faces, such as aluminum and copper, with carbon, nickel, titanium, silver, etc. can be used. As a charge collector for negative electrodes, besides copper, stainless steel, nickel, aluminum, titanium, a baked carbon, a conductive polymer, electrically conductive glass, an aluminum-Cd alloy, etc., it is the purpose of an adhesive property, conductivity, and oxidation-resistant improvement, and what processed front faces, such as copper, with carbon, nickel, titanium, silver, etc. can be used. About such material, it is also possible to oxidize a front face. About these configurations, the shape of the shape of others and a film and a sheet, the letter of a network, punch or the thing by which expanded one was carried out, a lath object, a porosity object, a foam, the organizer of a fiber group, etc. are used. [shape / of foil] Although especially limitation is thin, a 1-500-micrometer thing is used.

[0015] It is Li1-z to the main constituent of a positive active material. When using [Mn2-x-y Mx Liy O4], it is also possible to add other positive active materials. for example, MnO2, MoO3, V2 O5, Lix CoO2, and Lix NiO2 etc. -a metallic oxide, and TiS2, MoS2 and NbSe3 etc. -- various kinds of matter in which absorption/emission is possible can be used for alkali-metal ion, such as intercalated graphite, such as a metal chalcogen ghost, the poly acene, poly para-phenylene, polypyrrole, the poly aniline, and disulfide, and a conductive polymer, and an anion Applying and carrying out the coat of the technology, such as electrolytic polymerization, plating, sintering, a mechano fusion, and vacuum evaporationo, as the addition method besides only mixing is mentioned. [0016] Moreover, it is desirable to be able to use the organic electrolytic solution, a solid polymer electrolyte, an inorganic solid electrolyte, fused salt, etc., for example, and to use the organic electrolytic solution also in this as an electrolyte. As an organic solvent of this organic electrolytic solution, propylene carbonate, ethylene carbonate, Butylene carbonate, diethyl carbonate, dimethyl carbonate, Ester, such as methylethyl carbonate and gammabutyrolactone Substitution tetrahydrofurans, such as a tetrahydrofuran and 2-methyl tetrahydrofuran, A dioxolane, diethylether, dimethoxyethane, diethoxy ethane, Ether, such as methoxyethoxy ethane, dimethyl sulfoxide, a sulfolane, a methyl sulfolane, an acetonitrile, methyl formate, methyl acetate. N-methyl pyrrolidone, a dimethyl formamide, etc. are mentioned, and these can be used as independent or a mixed solvent. moreover -- as a supporting-electrolyte salt -- LiClO4, LiPF6, LiBF4, LiAsF6, LiCF3 SO3, LiN (CF3 SO2)2, LiN (C2 F5 SO2)2, and LiN (CF3 SO2) (C4 F9 SO2)2 etc. -- it is mentioned On the other hand, what melted the above supportingelectrolyte salts as a solid polymer electrolyte into polymer, such as a polyethylene oxide, the bridge formation object and poly force FAZEN, and its bridge formation object, can be used. Furthermore, inorganic solid electrolytes, such as Li3 N and LiI, are also usable. That is, what is necessary is just nonaqueous electrolyte of

lithium ion conductivity.

[0017] As separator, the transmittance of ion is excellent and an insulating thin film with a mechanical strength can be used. The sheet built with the polymer of olefin systems, such as organic-solvent-proof nature, hydrophobic shell polypropylene, and polyethylene, the glass fiber, the polyvinylidene fluoride, the polytetrafluoroethylene, etc., a micropore film, a nonwoven fabric, and cloth are used. The aperture of separator is a thing of the range generally used for a cell, for example, is 0.01-10 micrometers. Moreover, the same is said of the thickness, and it is the thing of the range generally used for a cell, for example, is 5-300 micrometers.

[0018] As a reason a charge-and-discharge property, especially whose cycle property improve, although it is not necessarily clear, it is considered as follows. Generally, in the interior of a cell, the various impurities which do not participate in the charge and discharge of a cell are included in many cases. For example, LiPF6 When using for an electrolyte, the salt itself can carry in an impurity or it is possible to produce HF (fluoric acid) reacting with the water of the ultralow volume contained in the interior of a cell, or a solvent. Although an ion conductivity high coat like a lithium carbonate is formed between the electrolytic solution and a carbon material on a carbon-material front face in the case of lithium occlusion, if an acid like fluoric acid exists after the time of this coat formation, or formation, an ion conductivity low lithium halide will be produced. The lithium halide produced in the interface of a carbon material and the electrolytic solution bars occlusion discharge of a lithium, as a result, increases the interfacial resistance of a negative electrode, and is considered to be one of the causes which service capacity reduces. Then, I thought that this problem would be solvable by reducing the amount of the fluoric acid which exists in the interior of a cell. That is, by the manganic acid lithium which is a positive active material thinking that decomposition will be promoted in catalyst, and using the manganic acid lithium which has the Spinel structure which replaced some manganese by elements other than a lithium and manganese, although it thinks that it is generated by the reaction of an electrolyte and water, this fluoric acid is ****, when the fluoric acid which drops the catalyst-activity exerted on decomposition etc. and is produced by the decomposition reaction of water and an electrolyte is reduced. Furthermore, this catalyst-activity is LiMn 2O4 which has Spinel structure. Gamma-MnO 2 which is the matter after charge By being the highest and replacing some manganese by elements other than a lithium and manganese This gamma-MnO 2 By using the manganic acid lithium which has the Spinel structure which suppressed generation, thought that catalytic activity could be reduced and replaced some manganese by elements other than a lithium and manganese Since interfacial-resistance increase of the carbon material of a negative electrode was suppressed and the cycle property improved, it resulted in this invention. Moreover, it turns out that its stability of an active material own [hot] improves catalytic activity not only reduced the manganic acid lithium which has the Spinel structure which replaced a part of this manganese by elements other than a lithium and manganese, but, and a hot cycle property also improves.

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EXAMPLE

[Example] Hereafter, this invention is explained based on an example.

[0020] (Example 1) The acetic-acid lithium dihydrate, manganese acetate (II)4 hydrate, and the calcium-acetate monohydrate were mixed so that Li and the ratio of Mn and calcium might be set to 1.10:1.95:0.05, and this was dissolved in the acetic acid. It agitated applying heat, the acetic acid was evaporated from the solution which dissolved completely, and the mixed salt was obtained. Temporary baking of this mixed salt was carried out at 500 degrees C, and it calcinated at 850 degrees C among air. As a result of grinding the obtained baking object and performing this XRD measurement, it turns out that the manganic acid lithium which has Spinel structure is obtained. Next, when the distributed state of calcium was observed by energy-dispersion type electron probe microanalysis (EPMA), calcium was distributed all over the manganic acid lithium. Let this powder be Powder A. [0021] (Example 2) The acetic-acid lithium dihydrate, manganese acetate (II)4 hydrate, and the chromium acetate (III) were mixed so that the ratio of Li, and Mn and Cr might be set to 1.10:1.95:0.05, and this was dissolved in the acetic acid. It agitated applying heat, the acetic acid was evaporated from the solution which dissolved completely, and the mixed salt was obtained. Temporary baking of this mixed salt was carried out at 500 degrees C, and it calcinated at 850 degrees C among air. As a result of grinding the obtained baking object and performing this XRD measurement, it turns out that the manganic acid lithium which has Spinel structure is obtained. Next, when the distributed state of chromium was observed by EPMA, chromium was distributed all over the manganic acid lithium. Let this powder be Powder B.

[0022] (Example 3) The acetic-acid lithium dihydrate, manganese acetate (II)4 hydrate, and the acetic-acid indium (III) hydrate were mixed so that the ratio of Li, and Mn and In might be set to 1.10:1.95:0.05, and this was dissolved in the acetic acid. It agitated applying heat, the acetic acid was evaporated from the solution which dissolved completely, and the mixed salt was obtained. Temporary baking of this mixed salt was carried out at 500 degrees C, and it calcinated at 850 degrees C among air. As a result of grinding the obtained baking object and performing this XRD measurement, it turns out that the manganic acid lithium which has Spinel structure is obtained. Next, when the distributed state of an indium was observed by EPMA, the indium was distributed all over the manganic acid lithium. Let this powder be Powder C.

[0023] (Example 4) The acetic-acid lithium dihydrate, manganese acetate (II)4 hydrate, and acetic-acid terbium (III) 4 hydrate were mixed so that Li and the ratio of Mn and Tb might be set to 1.10:1.95:0.05, and this was dissolved in the acetic acid. It agitated applying heat, the acetic acid was evaporated from the solution which dissolved completely, and the mixed salt was obtained. Temporary baking of this mixed salt was carried out at 500 degrees C, and it calcinated at 850 degrees C among air. As a result of grinding the obtained baking object and performing this XRD measurement, it turns out that the manganic acid lithium which has Spinel structure is obtained. Next, when the distributed state of a terbium was observed by EPMA, the terbium was distributed all over the manganic acid lithium. Let this powder be Powder D.

[0024] (Example 5) The acetic-acid lithium dihydrate, manganese acetate (II)4 hydrate, and magnesium-acetate (II)4 hydrate were mixed so that Li and the ratio of Mn and Mg might be set to 1.10:1.95:0.05, and this was dissolved in the acetic acid. It agitated applying heat, the acetic acid was evaporated from the solution which dissolved completely, and the mixed salt was obtained. Temporary baking of this mixed salt was carried out at 500 degrees C, and it calcinated at 850 degrees C among air. As a result of grinding the obtained baking object and performing this XRD measurement, it turns out that the manganic acid lithium which has Spinel structure is obtained. Next, when the distributed state of a chromium element was observed by EPMA, magnesium was distributed all over the manganic acid lithium. Let this powder be Powder E.

[0025] (Example 6) The coin type nonaqueous electrolyte cell shown in <u>drawing 1</u> as follows was made as an experiment, using the powder A, B, C, D, and E obtained in the above-mentioned example 1 as a positive active material. The positive electrode 1 mixed Powder A - Powder E, acetylene black, and polytetrafluoroethylene powder by the weight ratio 85:10:5, added toluene, and kneaded it enough. This was fabricated with a thickness of 0.8mm in the shape of a sheet with the roller press. next, this -- the diameter of 16mm -- it pierced circularly, and

dried at 200 degrees C under reduced pressure for 15 hours, and the positive electrode 1 was obtained The positive electrode 1 was stuck by pressure and used for the positive-electrode can 4 to which the positive-electrode charge collector 6 was attached. As a negative-electrode active material, it is an artificial graphite (the size (Lc) of the crystal of the spacing (d002) by 6 micrometers of mean particle diameters and the X-ray diffraction method of C shaft orientations by 3.37A mixed 550A and polytetrafluoroethylene powder by the weight ratio 95:5, added toluene, and kneaded enough.). This was fabricated with a thickness of 0.1mm in the shape of a sheet with the roller press. next, this -- the diameter of 16mm -- it pierced circularly, it dried at 200 degrees C under reduced pressure for 15 hours, and the negative electrode 2 was obtained The negative electrode 2 was stuck by pressure and used for the negative-electrode can 5 to which the negative-electrode charge collector 7 was attached. It is LiPF6 to the partially aromatic solvent of the volume ratio 1:1 of ethylene carbonate and diethyl carbonate. The fine porous membrane made from polypropylene was used for separator 3 using the electrolytic solution which carried out 1 mol/l dissolution. The coin type lithium cell with a diameter [of 20mm] and a thickness of 1.6mm was produced using the above-mentioned positive electrode, a negative electrode, the electrolytic solution, and separator. Let the cells using this powder A-E be a cell (A) - a cell (E), respectively.

[0026] (Example of comparison) The acetic-acid lithium dihydrate and manganese acetate (II)4 hydrate were mixed so that the ratio of Li and Mn might be set to 1.10:2.00, and this was dissolved in the acetic acid. It agitated applying heat, the acetic acid was evaporated from the solution which dissolved completely, and the mixed salt was obtained. Temporary baking of this mixed salt was carried out at 500 degrees C, and it calcinated at 850 degrees C among air. As a result of grinding the obtained baking object and performing this XRD measurement, it turns out that the manganic acid lithium which has Spinel structure is obtained. Let this powder be Powder C. Instead of Powder A, Powder F was used and the cell was produced like the example 3 except it. Let the obtained cell be a comparison cell (F).

[0027] These cells (A) The charge and discharge test was performed using - (E) and the comparison cell (F). Charge final voltage was set to 4.2V, the discharge final voltage was set to 3.0V, and 1mA and the test temperature performed constant-current charge and discharge for the charge and discharge current at 20 degrees C and 40 degrees C. The result of the service capacity of obtained 5 cycle eye was shown in Table 1. Moreover, the number of cycles at the time of service capacity falling to 60% of the first stage as a cycle life was measured.

[Table 1]

Table I				
瑶 池	20℃での 放電容量 (mAh)	サイクル寿命 (サイクル数)	40℃での 放電容量 (mAh)	サイクル寿命 (サイクル数)
電池A	1 6	285	1 7	1 4 5
電池B	1 6	2 6 5	1 7	1 4 0
電池C	1 6	3 0 0	1 7	1 6 0
電池D	1 6	265 .	1 7	1 4 0
電池E	1 6	275	1 7	1 4 5
比較電池F	1 7	8 0	1 4	1 0

[0029] Although the cell (A) - cell (E), and comparison cell (F) using Powder A - Powder E were compared and early service capacity did not change, a cycle life is good and a bird clapper is known. Moreover, in the elevated temperature of 40 degrees C, it turns out that early service capacity increases and a cycle life also improves. That is, there is little electrolytic disassembly by using the positive active material of this invention, and since generation of fluoric acid is suppressed by this, in the surface lining generated on a carbon front face, it is possible [it] that the coat of the component like the low lithium carbonate of comparatively resistance or a lithium oxide in which not the high lithium fluoride of the resistance generated under existence of fluoric acid but a fluorine does not participate is formed, interfacial-resistance increase is suppressed, and a cycle property improves. moreover, since the 40-degree

C hot cycle property is also excellent, not only in electrolytic disassembly In the above-mentioned example which can consider that elution of the manganese to the own electrolytic solution of a positive active material is also suppressed, and capacity deterioration of an active material is also suppressed To the main constituent in a positive active material, Li [Mn1.95calcium0.05Li 0.10O4], Li [Mn1.95Cr0.05Li 0.10O4] and Li [Mn1.95In0.05Li 0.10O4], Li [Mn1.95Tb0.05Li 0.10O4] Although Li [Mn1.95Mg0.05Li 0.10O4] was mentioned about the lithium secondary battery using the artificial graphite as a negative-electrode material, the same effect was checked about other substitution elements and negative-electrode material. In addition, this invention is not limited to the start raw material, the manufacture method, a positive electrode, a negative electrode, an electrolyte, separator, a cell configuration, etc. of the active material indicated by the above-mentioned example.

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DESCRIPTION OF DRAWINGS

[Brief Description of the Drawings]

[Drawing 1] It is the cross section of the coin type nonaqueous electrolyte cell concerning the example of this invention.

[Description of Notations]

- 1 Positive Electrode
- 2 Negative Electrode
- 3 Separator
- 4 Positive-Electrode Can
- 5 Negative-Electrode Can
- 6 Positive-Electrode Charge Collector
- 7 Negative-Electrode Charge Collector

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DRAWINGS

